In the Claims

- (Currently amended) A natriuretic compound conjugate comprising:
 - (a) a biologically active natriuretic compound comprising:
 - (i) a natriuretic molecule NPR-A binding site; and
 - (ii) at least one modifying moiety conjugation site

wherein the <u>biologically active</u> natriuretic compound <u>is</u> eemprises a peptide or a biologically active peptide segment of a brain natriuretic peptide, atrial natriuretic peptide, C-type natriuretic peptide, [[or]] dendroaspis natriuretic peptide or a biologically active segment thereof; and

(b) at least one a modifying moiety attached to said modifying moiety conjugation site, wherein the modifying moiety has a formula:

each C is independently selected and is an alkyl moiety having m carbons and m is from 1 to 20; and

each PAG is independently selected and is a polyalkylene glycol moiety having n subunits and n is from 2 to 25;

each X is independently selected and is a linking moiety selected from the group consisting of -C-, -O-, -C(O)-, -NH-, -NHC(O)-, and -C(O)NH-, and o is from 1 to 15, and

wherein said natriuretic compound conjugate exhibits one or more advantages selected from the group consisting of increased resistance to enzymatic degradation relative to a corresponding unconjugated natriuretic compound, increased circulating half life, increased bioavailability, and prolonged duration of effect.

- (Original) The natriuretic compound conjugate of claim 1 further defined as retaining a
 therapeutically significant percentage of cGMP stimulating activity relative to the corresponding
 unconjugated natriuretic compound.
- (Original) The natriuretic compound conjugate of claim 1 further defined as retaining at least 30% of the cGMP stimulating activity of the corresponding unconjugated natriuretic compound.
- (Original) The natriuretic compound conjugate of claim 1 further defined as retaining at least 50% of the cGMP stimulating activity of the corresponding unconjugated natriuretic compound.
- (Previously presented) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound is hBNP.
- (Original) The natriuretic compound conjugate of claim 1 further defined as retaining at least 90% of the cGMP stimulating activity of the corresponding unconjugated natriuretic compound.
- (Original) The natriuretic compound conjugate of claim 1 further defined as more hydrophilic than a corresponding unconjugated natriuretic compound.
- (Original) The natriuretic compound conjugate of claim 1 further defined as more amphiphilic than a corresponding unconjugated natriuretic compound.
- (Original) The natriuretic compound conjugate of claim 1 further defined as more lipophilic than
 a corresponding unconjugated natriuretic compound.
- 10. (Cancelled)
- (Original) The natriuretic compound conjugate of claim 1 further defined as more resistant to
 protease degradation than a corresponding unconjugated natriuretic compound.
- 12. (Withdrawn) The natrituretic compound conjugate of claim 1 wherein the natrituretic compound comprises a sequence:

A¹PX¹MVQGSGCFGRX²MDRISSSSGLGCX³VLR (SEQ ID NO. 116).

wherein

A1 is an amino acid or series of amino acids native to a natriuretic peptide,

 X^1 , X^2 and X^3 are independently selected from the group consisting of Lys, Arg and Gly, and at least one of X^1 , X^2 and X^3 is a Lys.

- 13. (Cancelled)
- 14. (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound comprises:
 - (a) an amino acid sequence

wherein

 \mathbf{X}^1 is optionally present and when present is an amino acid sequence having from 1–10 amino acids;

 \mathbf{X}^3 is optionally present and when present is an amino acid sequence having from $1{\text -}10$ amino acids.

- (b) a disulfide bond between C^1 and C^2 to form a loop.
- 15. (Withdrawn) The natriuretic compound conjugate of claim 14 wherein X¹ is Arg or Gly.
- (Withdrawn) The natriuretic compound conjugate of claim 14 wherein X¹ is selected from the group consisting of:
 - (a) Lvs:
 - (b) Gly;
 - (c) Arg;
 - (d) SG-, GSG-, QGSG- (SEQ ID NO. 118), VQGSG- (SEQ ID NO. 119), MVQGSG- (SEQ ID NO. 120), PKMVQGSG- (SEQ ID NO. 121), and SPKMVQGSG- (SEQ ID NO. 122);

- hBNP segments of (d) comprising a substitution selected from the group consisting of Lys-to-Gly and Lys-to-Arg;
- (f) hBNP segments of (d) comprising a substitution selected from the group consisting of Gly-to-Lys and Arg-to-Lys;
- (g) hBNP segments of (d) comprising an inserted Lys;
- (h) N-terminal tails and C-terminal segments of N-terminal tails of natriuretic peptides;
- N-terminal tails and C-terminal segments of (h) comprising a substitution selected from the group consisting of Lys-to-Gly and Lys-to-Arg;
- N-terminal tails and C-terminal segments of (h) comprising a substitution selected from the group consisting of Gly-to-Lys and Arg-to-Lys;
- (k) N-terminal tails and C-terminal segments of (h) comprising an inserted Lys.
- 17. (Withdrawn) The natriuretic compound conjugate of claim 14 wherein X³ is selected from the group consisting of:
 - (a) Lys;
 - (b) Glv:
 - (c) Arg;
 - (d) hBNP segments KV, KVL, KVLR (SEQ ID NO. 107), KVLRR (SEQ ID NO. 106), and KVLRRH (SEO ID NO. 105); and
 - hBNP segments of (d) comprising a substitution selected from the group consisting of Lys-to-Gly and Lys-to-Arg;
 - hBNP segments of (d) comprising a substitution selected from the group consisting of Gly-to-Lys and Arg-to-Lys;
 - (g) hBNP segments of (d) comprising an inserted Lys;
 - (h) C-terminal tails and N-terminal segments of C-terminal tails of natriuretic peptides;

- C-terminal tails and N-terminal segments of C-terminal tails of (h) comprising a substitution selected from the group consisting of Lys-to-Gly and Lys-to-Arg;
- C-terminal tails and N-terminal segments of C-terminal tails of (h) comprising a substitution selected from the group consisting of Gly-to-Lys and Arg-to-Lys;
- (k) C-terminal tails and N-terminal segments of C-terminal tails of (h) comprising an inserted Lys.
- 18. (Withdrawn) The natriuretic compound conjugate of claim 14 wherein the natriuretic compound comprises a sequence selected from the group consisting of:
 - (a) SPKMVQGSGCFGRKMDRISSSSGLGCKVL (SEQ ID NO. 123);
 - (b) SPKMVOGSGCFGRKMDRISSSSGLGC (SEQ ID NO. 124); and
 - segments (a) or (b) comprising a substitution selected from the group consisting of Lysto-Gly and Lys-to-Arg.
- (Withdrawn) The natriuretic compound conjugate of claim 14 wherein X¹ comprises a 1-9 amino acid residue sequence from the N-terminus of hBNP.
- (Withdrawn) The natriuretic compound conjugate of claim 14 wherein X¹ comprises SPX³MVQGSG (SEQ ID NO: 125), and wherein X² comprises a modifying moiety conjugation site.
- (Withdrawn) The natriuretic compound conjugate of claim 14 wherein X³ comprises a 1-6 amino acid residue sequence from the C-terminus of hBNP.
- (Withdrawn) The natriuretic compound conjugate of claim 14 wherein X³ comprises KVLRRH (SEQ. ID. NO: 105), KVLRR (SEQ ID NO. 106), KVLR (SEQ ID NO. 107), KVL, KV or K.
- (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound comprises a native hBNP sequence (SEQ ID NO. 73) having one or more mutations selected from the group consisting of Lys3Arg, Lys14Arg, Arg30Lys, Lys27Arg, and Arg31Lys.
- (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound comprises a native hBNP sequence (SEO ID NO. 73), having one or more insertions or deletions.

- (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound comprises a native hBNP amino acid sequence (SEQ ID NO. 73) and a N-terminal or C-terminal Lvs.
- (Withdrawn) The natriuretic compound conjugate of claim 1 further defined as:
 - (a) comprising a multipeptide comprising two or more amino acid sequences encoding a natriuretic compound;
 - optionally comprising a spacer sequence between each set or adjacent natriuretic compound encoding sequences;
 - (c) optionally comprising an extension at either or both ends of the multipeptide, the extension comprising one or more amino acids.
- (Withdrawn) The natriuretic compound conjugate of claim 26 wherein the natriuretic peptide
 units each comprise hBNP (SEQ ID NO. 73) or a biologically active analog, segment or segment
 analog thereof.
- (Original) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound consists of a native BNP.
- (Original) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound consists of a native hBNP.
- (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound consists of a native ANP.
- (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound consists of a canine BNP.
- (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound consists of urodilatin.
- (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound consists of DNP.

 (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound comprises an amino acid sequence:

X¹MVQGSGCFGRX²MDRISSSSGLGCX³ (SEQ ID NO. 126),

wherein X^1 , X^2 and X^3 are each independently selected from the group consisting of Lys, Gly and Arg, with the proviso that at least one of X^1 , X^2 and X^3 is Arg or Gly.

- 35. (Withdrawn) The natriuretic compound conjugate of claim 34 wherein the sequence comprises:
 - (a) N-terminal to X¹, an extension selected from the group consisting of: SPK, PK and K;
 and/or
 - (b) C-terminal to X³, an extension selected from the group consisting of -VLRRH (SEQ ID NO: 19), -VLRR (SEQ ID NO: 20), -VLR, -VL, and -V.
- (Withdrawn) The natriuretic compound conjugate of claim 34 wherein X¹ is Lys, X² is Arg and X³ is Arg.
- (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound comprises an amino acid sequence:

CFGRX1MDRISSSSGLGCX2 (SEO ID NO: 21).

wherein X¹ and/or X² comprises a modifying moiety conjugation site coupled to the modifying moiety.

- (Withdrawn) The natriuretic compound conjugate of claim 37 wherein X¹ comprises Lys coupled to the modifying moiety.
- (Withdrawn) The natriuretic compound conjugate of claim 37 wherein X² comprises Lys coupled to the modifying moiety.
- 40. (Original) The natriuretic compound conjugate of claim 1 wherein the modifying moiety conjugation site comprises a moiety selected from the group consisting of natural or non-natural amino acid side chains, an N-terminus of the natriuretic compound, and a C-terminus of the natriuretic compound.

- (Original) The natriuretic compound conjugate of claim 40 wherein the modifying moiety conjugation site is a Lys side chain.
- (Original) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound conjugate includes only one modifying moiety.
- 43. (Withdrawn) The natriuretic compound conjugate of claim 1 wherein:
 - the natriuretic compound comprises a Lys³ to Cys²⁶ segment of hBNP (SEQ ID NO. 127) and a disulfide bond coupling Cys¹⁰ of the segment to the Cys²⁶;
 - a single modifying moiety coupled to the natriuretic compound at the Lys³, wherein the amino acid sequence of hBNP is SEQ ID NO. 73.
- 44. (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound comprises a Cys¹⁰ to Cys²⁶ segment of hBNP (SEQ ID NO. 128) and a disulfide bond coupling the Cys¹⁰ to the Cys²⁶, wherein said natriuretic compound is a monoconjugate including a single modifying moiety coupled thereto at Lys¹⁴ of the segment.
- 45. (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound comprises a Cys¹⁰ to Lys²⁷ segment of hBNP (SEQ ID NO. 129), wherein said natriuretic compound is a monoconjugate including a single modifying moiety coupled thereto at Lys²⁷ of the segment.
- 46. (Withdrawnl) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound comprises a Cys¹⁰ to His³² (SEQ ID NO. 130) segment of hBNP and a disulfide bond coupling the Cys¹⁰ to Cys²⁶ of the segment, wherein said natriuretic compound is a monoconjugate including a single modifying moiety coupled thereto at Lys²⁷ of the segment.
- 47. (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound comprises a Cys¹⁰ to Cys²⁶ segment of hBNP (SEQ ID NO. 128) and a disulfide bond coupling the Cys¹⁰ to the Cys²⁶; wherein the natriuretic compound is a monoconjugate including a single modifying moiety coupled thereto at the N-terminus of the natriuretic compound.
- 48. (Withdrawn) The natriuretic compound conjugate of claim 1 wherein:
 - (a) the natriuretic compound consists of the hBNP amino acid sequence; and

- (b) the natriuretic compound conjugate is a diconjugate comprising:
- (c) a modifying moiety coupled to the natriuretic peptide at Lys³ of the hBNP amino acid sequence, wherein the amino acid sequence of hBNP is SEO ID NO. 73, and.
- (d) a modifying moiety coupled to the natriuretic peptide at Lys¹⁴ of the hBNP amino acid sequence, wherein the amino acid sequence of hBNP is SEO ID NO. 73.
- 49. (Withdrawn)The natriuretic compound conjugate of claim 1 wherein:
 - (a) the natriuretic compound is hBN, wherein the amino acid sequence of hBNP is SEQ ID NO.73; and
 - (b) the natriuretic compound conjugate is a diconjugate comprising:
 - a modifying moiety coupled to the natriuretic peptide at Lys³ of the hBNP amino acid sequence; and
 - a modifying moiety coupled to the natriuretic peptide at Lys²⁷ of the hBNP amino acid sequence.
- 50. (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the natriuretic compound sequence comprises N-terminal tail and the modifying moiety is coupled to an amino acid which is positioned in the N-terminal tail.
- 51. (Withdrawn) The natriuretic compound conjugate of claim 50 wherein the N-terminal tail consists of a native sequence of an N-terminal tail of a natriuretic peptide or a C-terminal segment of an N-terminal tail of a natriuretic peptide.
- 52.-85. (Cancelled).
- 86. (Cancelled).
- 87. (Cancelled).
- 88. (Cancelled).
- 89. (Original) The natriuretic compound conjugate of claim 86 wherein the polyalkylene glycol moiety comprises a polyethylene glycol moiety.

- 90. (Original) The natriuretic compound conjugate of claim 86 wherein the polyalkylene glycol moiety has from 2 to 25 polyalkylene glycol subunits.
- 91. (Original) The natriuretic compound conjugate of claim 86 wherein the polyalkylene glycol moiety has from 2 to 20 polyalkylene glycol subunits.
- 92. (Original) The natriuretic compound conjugate of claim 86 wherein the polyalkylene glycol moiety has from 2 to 15 polyalkylene glycol subunits.
- 93. (Original) The natriuretic compound conjugate of claim 86 wherein the polyalkylene glycol moiety has from 2 to 10 polyalkylene glycol subunits.
- 94. (Original) The natriuretic compound conjugate of claim 86 wherein the modifying moiety further comprises a linear or branched alkyl moiety.
- 95. (Cancelled).
- 96. (Original) The natriuretic compound conjugate of claim 94 wherein the alkyl moiety has from 1 to 20 carbons.
- 97. (Original) The natriuretic compound conjugate of claim 94 wherein the alkyl moiety has from 1 to 18 carbons.
- 98. (Original) The natriuretic compound conjugate of claim 94 wherein the alkyl moiety has from 1 to 16 carbons.
- 99. (Cancelled)
- 100. (Original) The natriuretic compound conjugate of claim 94 wherein the modifying moiety renders the natriuretic compound conjugate more lipophilic than a corresponding unconjugated natriuretic compound.
- 101. (Original) The natriuretic compound conjugate of claim 94 wherein the modifying moiety comprises a bond coupling the polyalkalene glycol moiety to the alkyl moiety which bond is hydrolysable in vivo.
- 102. (Withdrawn) The natriuretic compound conjugate of claim 1 wherein the modifying moiety comprises a linear or branched polyalkylene glycol moiety coupled to the natriuretic compound and a

linear or branched alkyl moiety coupled to the polyalkalene glycol moiety at a site which is distal relative to the natriuretic compound.

- 103. (Original) The natriuretic compound conjugate of claim 1 wherein the modifying moiety comprises a linear or branched alkyl moiety coupled to the natriuretic compound and a polyalkylene glycol moiety coupled to the alkyl moiety at a site which is distal relative to the natriuretic compound.
- 104. (Cancelled)
- 105. (Original) The natriuretic compound conjugate of claim 1 wherein the modifying moiety is coupled to the natriuretic compound by a bond that is hydrolysable *in vivo*.
- 106. (Original) The natriuretic compound conjugate of claim 1 wherein the modifying moiety is coupled to the natriuretic compound by a bond that is hydrolysable in the bloodstream.
- 107. (Original) The natriuretic compound conjugate of claim 1 wherein the modifying moiety is coupled to the natriuretic compound by a bond that is not hydrolysable *in vivo*.
- 108. (Original) The natriuretic compound conjugate of claim 1 wherein the modifying moiety is coupled to the natriuretic compound by a bond that is not hydrolysable in the bloodstream.
- 109. (Original) The natriuretic compound conjugate of claim 1 wherein the modifying moiety is coupled to the natriuretic compound by a bond selected from the group consisting of ester, carbonate, carbamate, amide, ether, and amine.
- 110. (Original) The natriuretic compound conjugate of claim 1 wherein the modifying moiety is hydrolysable in vivo to yield a pegylated natriuretic compound.
- 111. (Currently amended) The natriuretic compound conjugate of claim 110 wherein the modifying moiety is hydrolysable in vivo to yield a pegylated natriuretic compound comprising one or more PEG moieties having from ±2 to 6 PEG units.
- (Original) A pharmaceutical formulation comprising the natriuretic compound conjugate of claim
- 113. (Original) The pharmaceutical formulation of claim 112 formulated for a route of delivery selected from the group consisting of enteral, perenteral, oral, subcutaneous, sublingual, buccal, nasal, intravenous and intramuscular.

- 114. (Withdrawn) A method of treating a condition characterized by an excessive level of extracellular fluid, the method comprising administering to a subject in need thereof a pharmaceutically acceptable amount of a natriuretic compound conjugate of claim 1.
- 115. (Withdrawn) The method of claim 114 wherein the condition comprises congestive heart failure.
- 116. (Withdrawn) The method of claim 114 wherein the condition comprises chronic congestive heart failure
- 117. (Withdrawn) The method of claim 114 wherein the condition comprises acute congestive heart failure.
- 118. (Withdrawn) The method of claim 114 wherein the natriuretic compound conjugate is self-administered.
- 119. (Withdrawn) The method of claim 114 wherein the natriuretic compound conjugate is orally administered.
- 120. (Withdrawn) The method of claim 114 wherein the natriuretic compound conjugate is administered via a route of administration selected from the group consisting of enteral, perenteral, oral, subcutaneous, sublingual, buccal, nasal, intravenous and intramuscular.
- 121. (Withdrawn) The method of claim 114 wherein the condition is hypertension.
- 122. (Withdrawn and currently amended) A method of making the natriuretic compound conjugate of claim 1, the method comprising:
 - a. conjugating a natriuretic peptide multipeptide comprising two or more natriuretic compound units wherein the natriuretic peptide is selected from the group consisting of a brain natriuretic peptide, atrial natriuretic peptide, C-type natriuretic peptide, dendroaspis natriuretic peptide and a biologically active segment thereof;
 - b. cleaving the natriuretic peptide multipeptide to yield natriuretic compound conjugate;
 - oxidizing the cleaved natriuretic compound conjugate to form one or more disulfide bonds in the natriuretic compound conjugate.

- 123. (Withdrawn) The method of claim 122 wherein the natriuretic compound comprises Cys¹⁰ to Cys²⁶ of hBNP (SEQ ID NO. 128) and step 122.c yields a disulfide bond between the Cys¹⁰ and Cys²⁶.
- 124. (Withdrawn) A method of making the natriuretic compound conjugate of claim 1, the method comprising:
 - making a multi-peptide natriuretic compound comprising two or more natriuretic compound units:
 - b. cleaving the natriuretic peptide multipeptide to yield natriuretic peptide compound;
 - c. conjugating the natriuretic compound to yield natriuretic compound conjugate;
 - d. oxidizing the cleaved natriuretic compound conjugate to form one or more disulfide bonds in the natriuretic compound conjugate.
- 125. (Withdrawn and currently amended) The method of claim 124 wherein the natriuretic compound comprises Cys¹⁰ to Cys²⁶ of hBNP (SEQ ID NO. 128) and step 124(c) 122.e yields a disulfide bond between the Cys¹⁰ and Cys²⁶.
- 126. (Withdrawn and currently amended) A method of making the natriuretic compound conjugate of claim 1, the method comprising:
 - a. making a multi-peptide natriuretic compound comprising two or more natriuretic compound units selected from the group consisting of brain natriuretic peptide, atrial natriuretic peptide, C-type natriuretic peptide, dendroaspis natriuretic peptide and a biologically active segment thereof;
 - b. cleaving the natriuretic peptide multipeptide to yield natriuretic compound;
 - oxidizing the cleaved natriuretic compound to form one or more disulfide bonds in the natriuretic compound; and
 - d. conjugating the natriuretic compound to the modifying moiety of claim 1.
- (Withdrawn) A modified pro-polynatriuretic peptide conjugate comprising:
 - at least one natureteic peptide unit having a modifying moiety conjugation site and an NPR-A binding site;

- at least one modifying moeity attached to the modifying moiety conjugation site of at least one of the natriuretic peptide units;
- c. a leader sequence; and
- d. an enzymatically cleavable spacer coupling the leader sequence to a first natriuretic peptide conjugate.
- 128. (Withdrawn) A natriuretic peptide analog comprising an amino acid sequence having at least one modifying moiety conjugation site, an NPR-A binding region and at least one substituted Lys residue therein as compared to a native natriuretic peptide amino acid sequence, wherein said substituted Lys residue is not the amino acid modifying moiety conjugation site.
- 129. (Withdrawn) The natriuretic peptide analog of claim 128, wherein the native natriuretic peptide has the amino acid sequence SEQ ID NO. 73, wherein the one or more substituted Lys residues comprise a substitution selected from the group consisting of: Lys3Gly, Lys3Arg, Lys14Gly, Lys14Arg, Lys27Gly, or Lys27Arg.
- 130. (Withdrawn) The natriuretic peptide analog of claim 128 comprising a structure:

SPKMVOGSGCFGRX1MDRISSSSGLGCX2VLRRH (SEO ID NO: 131)

- a. wherein X^1 is Lys and X^2 is other than Lys, or X^1 is Lys and X^2 is other than Lys, or X^1 and X^2 are other than Lys.
- 131. (Withdrawn) The natriuretic peptide analog of claim 130 wherein X¹ is Lys and X² is Arg or Gly, or X¹ is Lys and X² is Arg or Gly, or X¹ and X² are independently selected and are Arg or Gly.
- 132. (Withdrawn) A natriuretic peptide analog comprising a structure:

CFGRX1MDRISSSSGX2GC (SEO ID NO: 132)

wherein X^1 is an amino acid that does not comprise a conjugation site, and X^2 is an amino acid that comprises a modifying moiety conjugation site.

- 133. (Withdrawn) The natriuretic peptide analog of claim 132 wherein X¹ is Arg and X² is Lys.
- 134. (Withdrawn) A natriuretic peptide analog having a structure:

X1-CFGRX3MDRISSSSGLGC-X2 (SEQ ID No. 117)

wherein X^1 is an amino acid sequence having from 1 to 10 amino acids, X^2 is an amino acid sequence having from 1 to 10 amino acids, and X^3 is other than Lys.

- 135. (Withdrawn) The natriuretic peptide analog of claim 134 wherein X3 is Arg or Gly.
- 136. (Withdrawn) The natriuretic peptide analog of claim 134 wherein X¹ is SPY¹MVQGSG (SEQ ID NO: 133), wherein Y¹ comprises a modifying moiety conjugation site.
- 137. (Withdrawn) The natriuretic peptide analog of claim 134 wherein X^{1} is selected from the group consisting of:
 - a. N-terminal tails and C-terminal segments of N-terminal tails of natriuretic peptides:
 - N-terminal tails and C-terminal segments of a comprising a substitution selected from the group consisting of Lys-to-Gly and Lys-to-Arg;
 - N-terminal tails and C-terminal segments of a comprising a substitution selected from the group consisting of Gly-to-Lys and Arg-to-Lys;
 - d. N-terminal tails and C-terminal segments of a comprising an inserted Lys.
- 138. (Withdrawn) The natriuretic peptide analog of claim 134 wherein X² is Y²VLRRH (SEQ. ID. NO: 134), wherein Y² is other than Lys.
- 139. (Withdrawn) The natriuretic peptide analog of claim 138 wherein Y² is Arg.
- 140. (Withdrawn) The natriuretic peptide analog of claim 134 wherein X² is selected from the group consisting of:
 - a. C-terminal tails and N-terminal segments of C-terminal tails of natriuretic peptides;
 - C-terminal tails and N-terminal segments of C-terminal tails of 137.a comprising a substitution selected from the group consisting of Lys-to-Gly and Lys-to-Arg;
 - C-terminal tails and N-terminal segments of C-terminal tails of 137.a comprising a substitution selected from the group consisting of Gly-to-Lys and Arg-to-Lys;

- d. C-terminal tails and N-terminal segments of C-terminal tails of 137.a comprising an inserted Lys.
- 141. (Withdrawn) A natriuretic peptide analog having a structure:

X1-CFGRX3MDRIGLGC-X2 (SEO ID No. 135)

wherein X^1 is a peptide of from 1 to 9 amino acids, X^2 is a peptide of from 1 to 6 amino acids, and X^3 is other than Lvs.

- 142. (Withdrawn) The natriuretic peptide analog of claim 140 wherein X³ is Arg or Gly.
- 143. (Withdrawn) The natriuretic peptide analog of claim 142 wherein X¹ is SPY¹MVQGSG (SEQ ID NO: 133), wherein Y¹ comprises a modifying moiety conjugation site.
- 144. (Withdrawn) The natriuretic peptide analog of claim 142 wherein X² is Y²VLRRH (SEQ. ID. NO: 134), wherein Y² is other than Lys.
- 145. (Withdrawn) The natriuretic peptide analog of claim 144 wherein Y² is Arg.
- 146. (Withdrawn) The natriuretic peptide analog of claim 144 wherein X³ is Arg, X¹ is a sequence SPKMVOGSG (SEO ID NO: 122) and X² is a sequence RVL.
- 147. (Withdrawn) A natriuretic peptide analog having a structure X^t-CFGRX³MDRIX^tGLGC-X² (SEQ ID NO. 136) wherein
 - a. X1 is an amino acid sequence of from 1 to 10 amino acids,
 - X² is an amino acid sequence of from 1 to 10 amino acids,
 - c. X4 is an amino acid sequence of from 1 to 4 amino acids; and
 - d. X3 is other than Lys.
- 148. (Withdrawn) The natriuretic peptide analog of claim 147 wherein neither X^1 nor X^2 is a sequence native to a natriuretic peptide.
- (Withdrawn The natriuretic peptide of claim 147 where X³ is Arg or Gly.

- 150. (Withdrawn) The natriuretic peptide of claim 147 where X1 is SPY1MVOGSG (SEO ID NO: 133) wherein Y1 comprises a modifying moiety conjugation site.
- (Withdrawn) The natriuretic peptide analog of claim 147 wherein X² is Y²VLRRH (SEQ. ID. 151. NO: 134), wherein Y2 is other than Lys.
- (Withdrawn) The natriuretic pentide analog of claim 151 wherein Y2 is Arg. 152.
- (Withdrawn) An hBNP analog comprising a substitution of Lys14Arg or Lys14Gly, wherein the 153 amino acid sequence of hBNP is SEO ID NO. 73.
- 154. (Withdrawn) An hBNP analog comprising a substitution of Lys27Arg or Lys27Gly, wherein the amino acid sequence of hBNP is SEO ID NO. 73.
- 155. (Withdrawn) An hBNP analog comprising a substitution of Lys3Arg or Lys3Gly, wherein the amino acid sequence of hBNP is SEQ ID NO. 73.
- 156. (Previously presented) A natriuretic compound conjugate comprising:
 - (a) a natriuretic compound comprising:

i.a natriuretic molecule NPR-A binding site; and

- ii. at least one modifying moiety conjugation site wherein the natriuretic compound comprises a peptide or a biologically active peptide segment of brain natriuretic peptide, atrial natriuretic peptide, C-type natriuretic peptide, or dendroaspis natriuretic peptide; and
- at least one modifying moiety attached to said modifying moiety conjugation site, (b) wherein the modifying moiety has a formula:

each C is independently selected and is an alkyl moiety having m carbons and m is from 1 to 20; and

each PAG is independently selected and is a polyalkylene glycol moiety having n subunits and n is from 2 to 25:

each X is independently selected and is a linking moiety selected from the group consisting of -C-, -O-, -C(O)-, -NH-, -NHC(O)-, and -C(O)NH-, and o is from 1 to 15, and

wherein said natriuretic compound retains a therapeutically significant percentage of cGMP stimulating activity relative to a corresponding unconjugated natriuretic compound.

- 157. (Previously presented) A natriuretic compound conjugate comprising:
 - (a) a natriuretic compound comprising:
 - i. a natriuretic molecule NPR-A binding site; and
 - ii. at least one modifying moiety conjugation site

wherein the natriuretic compound comprises a peptide or a biologically active peptide segment of brain natriuretic peptide, atrial natriuretic peptide, C-type natriuretic peptide, or dendroaspis natriuretic peptide; and

(b) at least one modifying moiety attached to said modifying moiety conjugation site, wherein the modifying moiety has a formula:

each C is independently selected and is an alkyl moiety having m carbons and m is from 1 to 20; and

each PAG is independently selected and is a polyalkylene glycol moiety having n subunits and n is from 2 to 25:

each X is independently selected and is a linking moiety selected from the group consisting of -C-, -O-, -C(O)-, -NH-, -NHC(O)-, and -C(O)NH-, and o is from 1 to 15, and

wherein said natriuretic compound conjugate retains at least 50% of the cGMP stimulating activity of a corresponding unconjugated natriuretic compound.

- 158. (Previously presented) A natriuretic compound conjugate comprising:
 - (a) a natriuretic compound comprising:
 - i. a natriuretic molecule NPR-A binding site; and
 - at least one modifying moiety conjugation site wherein the natriuretic compound comprises a peptide or a biologically active peptide segment of brain natriuretic peptide, atrial natriuretic peptide, C-type natriuretic peptide, or dendroaspis natriuretic peptide; and
 - (b) at least one modifying moiety attached to said modifying moiety conjugation site, wherein the modifying moiety has a formula:

each C is independently selected and is an alkyl moiety having m carbons and m is from 1 to 20; and

each PAG is independently selected and is a polyalkylene glycol moiety having n subunits and n is from 2 to 25:

each X is independently selected and is a linking moiety selected from the group consisting of -C-, -O-, -C(O)-, -NH-, -NHC(O)-, and -C(O)NH-, and o is from 1 to 15, and

wherein said natriuretic compound conjugate is more hydrophilic than a corresponding unconjugated natriuretic compound.

- (Previously presented) A natriuretic compound conjugate comprising:
 - (a) a natriuretic compound comprising:

- i. a natriuretic molecule NPR-A binding site; and
- at least one modifying moiety conjugation site wherein the natriuretic compound comprises a peptide or a biologically active peptide segment of brain natriuretic peptide, atrial natriuretic peptide, C-type natriuretic peptide, or dendroaspis natriuretic peptide; and
- (b) at least one modifying moiety attached to said modifying moiety conjugation site, wherein the modifying moiety has a formula:

each C is independently selected and is an alkyl moiety having m carbons and m is from 1 to 20; and

each PAG is independently selected and is a polyalkylene glycol moiety having n subunits and n is from 2 to 25:

each X is independently selected and is a linking moiety selected from the group consisting of -C-, -C(O)-, -NH-, -NHC(O)-, and -C(O)NH-, and o is from 1 to 15, and

wherein said natriuretic compound conjugate is more amphiphilic than a corresponding unconjugated natriuretic compound.

- 160. (Withdrawn) A natriuretic compound conjugate comprising:
 - a. a natriuretic compound comprising:
 - i. a natriuretic molecule NPR-A binding site; and
 - ii. at least one modifying moiety conjugation site; and
 - b. at least one modifying moiety attached to said modifying moiety conjugation site;

wherein the natriuretic compound conjugate is more lipophilic than a corresponding unconjugated natriuretic compound, wherein at least one modifying moiety does not consist of an alkyl moiety.

(Withdrawn) A compound having a formula:

wherein

each C is independently selected and is an alkyl moiety having m carbons and m is from 1 to 20; and

each PAG is independently selected and is a polyalkylene glycol moiety having n subunits and n is from 2 to 25;

each X is independently selected and is a linking moiety.

162. (Withdrawn) A compound having a formula:

$$C_m$$
-X-PAG_n PAG_m -X-C_m (Formula V)

wherein

each C is independently selected and is an alkyl moiety having m carbons and m is from 1 to 20; and

each PAG is independently selected and is a polyalkylene glycol moiety having n subunits and n is from 2 to 25:

each X is independently selected and is a linking moiety.

(Withdrawn) A compound having a formula;

PAG is a polyalkylene glycol moiety having n subunits and n is from 2 to 25;

X is O or N; and

each o is independently selected and is from 1 to 15.

164. (Withdrawn) A compound having a formula:

PAG is a polyalkylene glycol moiety having n subunits and n is from 2 to 25;

X is O or N; and

each o is independently selected and is from 1 to 15.

165. (Withdrawn) A compound having a formula:

$$C_{n}$$
- \times -PAG $_{n}$ PAG_{n} - $\times C_{m}$ (Formula VIII)

each C is independently selected and is an alkyl moiety having m carbons and m is from 1 to 20; and

each PAG is independently selected and is a polyalkylene glycol moiety having n subunits and n is from 2 to 25:

each X is independently selected and is a linking moiety.

is from 1 to 15

166. (Withdrawn) A compound having a formula;

each C is independently selected and is an alkyl moiety having m carbons and m is from 1 to 20; and

each PAG is independently selected and is a polyalkylene glycol moiety having n subunits and n is from 2 to 25;

each X is independently selected and is a linking moiety;

is from 1 to 15.

167. (Withdrawn) A method of making a compound of the formula:

wherein

each C is independently selected and is an alkyl moiety having m carbons and m is from 1 to 20; and

each PAG is independently selected and is a polyalkylene glycol moiety having n subunits and n is from 2 to 25:

each X is independently selected and is a linking mojety:

the method comprising:

reacting a compound of formula:

with a compound of formula:

$$X^2$$

where X^2 is a halide, and wherein the reaction is carried out in the presence of a base and a solvent to yield:

reacting the product of (a) with a compound of formula:

in the presense of a Lewis acid and a solvent to yield:

- 168. (Withdrawn) The method of claim 167 wherein the base is NaH and the solvent is tetrahydrofuran.
- 169. (Withdrawn) The method of claim 167 wherein the Lewis acid is BF3OEt2.
- 170. (Withdrawn) A method of making a compound of the formula:

$$\bigcap_{C_m-X-PAG_n}\bigcap_{PAG_m-X-C_m}\bigcap_{(Formula\ V)}$$

wherein

each C is independently selected and is an alkyl moiety having m carbons and m is from 1 to 20: and

each PAG is independently selected and is a polyalkylene glycol moiety having n subunits and n is from 2 to 25;

each X is independently selected and is a linking moiety;

the method comprising reacting the product of claim 161 with paranitrochloroformate or disuccimidyl carbonate.

171. (Withdrawn) A method of making a compound of the formula:

wherein

PAG is a polyalkylene glycol moiety having n subunits and n is from 2 to 25;

X is O or N; and

each o is independently selected and is from 1 to 15;

the method comprising:

reacting a compound of formula:

wherein o is as defined above, with a compound of formula:

HO-PAG_n-X

where X is -NH or -OH;

in solvent, to yield a compound of formula:

172. (Withdrawn) A method of making a compound of the formula:

(Formula VII)

wherein

PAG is a polyalkylene glycol moiety having n subunits and n is from 2 to 25;

X is O or N; and

each o is independently selected and is from 1 to 15;

the method comprising activating a product of claim 170 using an activating agent selected from the group consisting of disuccinimidyl carbonate, paranitrochloroformate, phosgene and N-hydroxysuccinimide.

173. (Withdrawn) A method of making a compound of the formula:

$$\bigcap_{\mathsf{C}_m - \mathsf{X} - \mathsf{PAG}_n} \bigcap_{\mathsf{PAG}_n - \mathsf{X} - \mathsf{C}_m} \mathsf{OH}$$

wherein

each C is independently selected and is an alkyl moiety having m carbons and m is from 1 to 20; and

each PAG is independently selected and is a polyalkylene glycol moiety having n subunits and n is from 2 to 25;

each X is independently selected and is a linking moiety.

is from 1 to 15:

the method comprising:

reacting the product of claim 161 with a compound of formula:

$$H_2N$$
 OH

in the presence of a base in a solvent.

- 174. (Withdrawn) The method of claim 173 wherein the base is K₂CO₃ and the solvent is an aqueous and/or organic solvent.
- 175. (Withdrawn) A method of making a compound of the formula:

$$C_{m}\text{-}\text{\times-$PAG}_{n} \qquad \qquad PAG_{n}\text{-}\text{\times-$C}_{m} \qquad \qquad (Formula~IX)$$

wherein

each C is independently selected and is an alkyl moiety having m carbons and m is from 1 to 20; and

each PAG is independently selected and is a polyalkylene glycol moiety having n subunits and n is from 2 to 25:

each X is independently selected and is a linking moiety;

is from 1 to 15:

the method comprising reacting a compound produced according to the method of claim 173 with N-hydroxysuccinimide.

176. (Withdrawn) A natriuretic peptide analog comprising a structure:

SPX1MMHX2SGCFGRRLDRIGSLSGLGCNVLRX3Y (SEQ ID NO. 137)

wherein X1 is Lvs. Arg or His, X2 is Lvs. Arg. His, and X3 is Arg or His.

- 177. (Withdrawn) The natriuretic peptide analog of claim 176 comprising a modifying moiety conjugated at the S residue.
- 178. (Withdrawn) A natriuretic peptide analog comprising a structure:

SPZ¹MVQGSG-CFGRZ²MDRISSSSX¹X²X³C (SEQ ID NO. 113)

wherein Z^1 is Arg or an amino acid other than Lys, and wherein Z^2 is Arg or an amino acid other than Lys, wherein X^1 is Gly, Met, Leu, Phe, Ile or a conservative substitution thereof, wherein X^2 is Leu, Trp, Tyr, Phe or a conservative substitution thereof, and wherein X^3 is Gly and Arg, or a conservative substitution thereof.

- 179. (Withdrawn) The natriuretic peptide analog of claim 178 where Z¹ is Lys and Z² is other than Lys.
- 180. (Withdrawn) A natriuretic peptide analog comprising a structure:

K CFKGKNDRX¹KX²QSGLX³C-NSFKY (SEQ ID NO. 114)

wherein X1 is T. a. R. H. P. T. E:

wherein X2 is K, N-methyl, Arg, S, D,P;

wherein X³ is Arg, K, Y, F, S, P, Orn, Har, Har, p-amidinophenyl Ala, I, any other amino acid that has a positive charge other than Gly, or Try.

181. (Withdrawn) The natriuretic peptide of claim 178 or 180 further defined as comprising a natriuretic peptide conjugate, comprising a modifying moiety conjugated to one or more of the Lys residues therein.